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(54) Title: TEST AND MODEL FOR INFLAMMATORY DISEASE

(57) Abstract: The present invention relates to a polynucleotide encoding the corneodesmosin protein having one or more nucleotide insertions, deletions or substitutions at one or novel positions. The invention also relates to the corneodesmosin protein having one or more amino acid insertions, deletions and substitutions. These nucleotide and amino acid polymorphisms are useful in diagnosing or determining susceptibility to corneodesmosin-mediated disease, for example inflammatory diseases including psoriasis, and in treating such disease. Host cells and transgenic non-human animals comprising polynucleotides or proteins of the invention are provided. Methods of screening for agents for use in treating corneodesmosin-mediated disease are also provided.

Test and Model for Inflammatory Disease

The present invention relates to nucleotide substitutions, deletions or insertions in the corneodesmosin gene, and the exploitation of these polymorphisms in the detection 5 and/or treatment of corneodesmosin mediated disease, for example inflammatory diseases including psoriasis. The present invention also relates to polynucleotides encoding the corneodesmosin protein, and having one or more nucleotide polymorphisms, and to a protein encoded by said polynucleotides. Also provided are transgenic non-human animals comprising the polynucleotides of the present 10 invention; and methods and kits for treating, diagnosing or determining susceptibility to corneodesmosin mediated disease, in particular by way of gene therapy.

In recent years, it has been recognised that there is considerable genetic diversity in 15 human populations, with common polymorphisms occurring on average at least every kilobase in the genome. Polymorphisms which affect gene expression or activity of the encoded gene product may account for susceptibility to, or expression of, disease conditions, either directly or through interaction with other genetic and environmental factors.

20 Understanding the molecular basis for disease, by sequencing the human genome and characterising polymorphisms, will enable the identification of those individuals at greatest risk of disease. This will allow the better matching of treatment and disease, and enable the production of new and improved targets for drugs. Screening and treatment of disease may also be better targeted to those in need, thus increasing the 25 cost-effectiveness of health-care provision.

One area in need of such approaches is the diagnosis and treatment of inflammatory diseases. Inflammation, which can be broadly defined as the destructive sequelae to activation of elements of the body's immune system, is a feature of many diseases

including infection, autoimmune disorders and benign and malignant hyperplasia. The identification of genetic factors which influence susceptibility to such disorders will provide important new insights into inflammatory disease, and may yield important new diagnostic and/or prognostic tests and treatments.

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Psoriasis is a chronic inflammatory cutaneous disorder which affects approximately 2% of the population in the UK and US. Psoriasis manifests itself as red scaly skin patches, principally on the scalp, elbows and knees, and is caused by epidermal hyperproliferation, and abnormal differentiation and infiltration of inflammatory cells.

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Psoriasis may also be associated with other inflammatory diseases such as arthritis, Crohn's disease, and HIV infection. Population, family, and twin studies all suggest an important genetic component in the pathogenesis of psoriasis, coupled with environmental triggers such as streptococcal infection and stress.

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Psoriasis is one of a number of autoimmune diseases that display significant human leukocyte antigen (HLA) associations. The analysis of population-specific HLA haplotypes has provided evidence that susceptibility to psoriasis is linked to the class I and II major histocompatibility complexes (MHC) on human chromosome 6 (Jenisch *et al.* (1998) *Am. J. Hum. Genet.* 63:191-199). These studies show that psoriasis consists of two distinct disease subtypes (Type I and Type II), which differ in age of onset and in the frequency of HLA types. Type I psoriasis has an age of onset of prior to 40 years and HLA types Cw6, B57, and DR7 are strongly increased. Patients with Type I psoriasis are much more likely to have a positive family history for the disease. In contrast, only about 10% of Cw6-positive individuals develop Type II psoriasis disease, with HLA-Cw2 being over-represented in this group.

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Linkage analysis and association studies suggest the presence of a major genetic determinant of psoriasis within the MHC, the strongest candidate gene marker being HLA-C. The most significant association has been shown between HLA-Cw6 and disease Type IA, which has the earliest onset of disease at 0 to 20 years. However,

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specific involvement of the HLA-Cw6 genotype in disease pathogenesis has yet to be established.

Recently, attention has focussed on non-HLA genes close to HLA-C, in particular the  
5 corneodesmosin gene (also known as the S gene), which is located approximately 160 kb telomeric of the HLA-C locus. The corneodesmosin gene consists of 2 exons spanning approximately 5.3 kb of genomic DNA sequence. Two corneodesmosin mRNAs of 2.2kb and 2.6kb, resulting from alternative splicing, have been described (Guerrin *et al.* (1998) *J. Biol. Chem.* **273**:22640-22647). Association studies (Ahnini  
10 *et al.* (1999) *Hum. Mol. Genet.* **8**:1135-1140) suggest a strong, significant association between a polymorphism at position 1243 of the corneodesmosin gene and psoriasis. A corneodesmosin gene haplotype was subsequently defined, which by TDT analysis was shown to have a strong, significant association with psoriasis (Allen *et al.* (1999) *Lancet* **353**:1589-90).

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In human epidermis and other cornified squamous epithelia, corneodesmosin is located in the desmosomes of the upper living layers, and in related structures of the cornified layers, the corneodesmosomes. During maturation of the cornified layers, the protein undergoes a series of cleavages, thought to be a prerequisite of  
20 desquamation (shedding of the cuticle or epidermis). Corneodesmosin is detected as a glycosylated and phosphorylated basic protein with an apparent molecular mass of 52-56 kDa. During stratum corneum maturation, corneodesmosin is progressively proteolysed until desquamation occurs. In superficial corneocytes, the 52-56 kDa form is no longer detected and immunoreactive fragments of 45 to 30 kDa predominate.  
25 Since location, biochemical characteristics and processing of corneodesmosin are similar in several mammals, it is likely that the protein is essential for the function of corneodesmosomes and corneocyte cohesion. It has been shown that expression of the 56kDa epidermal keratin polypeptide is increased in psoriatic lesions compared with normal skin and transformation of desmosomes into corneodesmosomes is altered in psoriatic epidermis.  
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Psoriasis affects approximately 6.4 million people in the US and causes varying ranges of physical discomfort, pain and disability. At present, the causes of psoriasis are unknown. There is no specific test for psoriasis or susceptibility thereto, and diagnosis 5 is based solely on clinical examination and skin histopathology.

It is likely that corneodesmosin is implicated in a range of skin diseases, including psoriasis. In this text, diseases in which corneodesmosin is implicated in the pathology will be referred to as "corneodesmosin-mediated disease".

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The present invention aims to overcome or ameliorate previous limitations in the art by providing means and methods for the detection and treatment of individuals having, or being susceptible to, corneodesmosin mediated disease, in particular inflammatory conditions such as psoriasis.

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In a first aspect, the present invention provides an isolated or recombinant polynucleotide comprising a nucleic acid sequence encoding the corneodesmosin gene of Figure 1, wherein said nucleic acid sequence comprises a nucleotide substitution, deletion or insertion at one or more of positions 6984, 7068, 7077, 7107, 7164, 8884, 20 8906, 8931, 9538, 9607, 9608, 9647, 9667, 9745, 9761, 9926, 9952, 9968, 10082, 10161, 10162, 10363, 11567, 11641, 11649, 11808, 11839, 11885, 11977, 12018, 12136, 12149; 12198, 12283, 12318, 12345, 12373, 12901, 13001, 13020, 13108, 13117, 13178, 13224, 13316, 13365, 13562, 13605, 13670, 13859, 13889 and 13914 of Figure 1. These novel polymorphisms in the corneodesmosin gene, at the positions indicated above, may be responsible for corneodesmosin mediated disease. In 25 particular, the polymorphisms of the present invention may be useful in identifying individuals susceptible or resistant to corneodesmosin-mediated disease, and in the diagnosis or treatment of such conditions. Preferred combinations of the polymorphisms of the invention are the haplotypes shown in Tables 10a and b. The 30 most preferred haplotype is B of Table 10a.

The polynucleotide of this invention is preferably DNA, or may be RNA or other options.

5 By "isolated" is meant a polynucleotide sequence which has been purified to a level sufficient to allow allelic discrimination. For example, an isolated sequence will be substantially free of any other DNA or protein product. Such isolated sequences may be obtained by PCR amplification, cloning techniques, or synthesis on a synthesiser. By recombinant is meant polynucleotides which have been recombined by the hand of  
10 man.

The corneodesmosin gene sequence shown in Figure 1 refers to the genomic clone of corneodesmosin, detailed in GenBank Accession No. AC006163 (a genomic clone of the MHC region on chromosome 6p21.3). The single nucleotide polymorphisms of  
15 the invention are shown in bold type and underlined on this figure, and have each been given a positional reference with respect to this figure. For reference and comparison with prior art publications, the positional references with respect to the coding sequence have also been given in Table 6, column 2, where nucleotide position 1 corresponds to the first nucleotide of exon 1 and nucleotides upstream of this are given  
20 a negative prefix.

A polymorphism is typically defined as two or more alternative sequences, or alleles, of a gene in a population. A polymorphic site is the location in the gene at which divergence in sequence occurs. Examples of the ways in which polymorphisms are manifested include restriction fragment length polymorphisms, variable number of tandem repeats, hypervariable regions, minisatellites, di- or multi-nucleotide repeats, insertion elements and nucleotide deletions, additions or substitutions. The first identified allele is usually referred to as the reference allele, or the wild type. Additional alleles are usually designated alternative or variant alleles. Herein, the sequence exactly as shown in Figure 1 is designated the reference sequence, and is not  
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part of the invention. Nucleic acid sequences of the present invention which differ from the sequence of Figure 1 at one or more of the positions indicated above may be referred to as variants of Figure 1.

- 5 A single nucleotide polymorphism is a variation in sequence between alleles at a site occupied by a single nucleotide residue. Single nucleotide polymorphisms (SNP's) arise from the substitution, deletion or insertion of a nucleotide residue at a polymorphic site. Typically, this results in the site of the variant sequence being occupied by any base other than the reference base. For example, where the reference  
10 sequence contains a "T" base at a polymorphic site, a variant may contain a "C", "G" or "A" at that site. Single nucleotide polymorphisms may result in corresponding changes to the amino acid sequence. For example, substitution of a nucleotide residue may change the codon, resulting in an amino acid change. Similarly, the deletion or insertion of three consecutive bases in the nucleic acid sequence may result in the  
15 insertion or deletion of an amino acid residue. For ease of reference, where a single nucleotide polymorphism of the present invention results in the insertion or deletion of a nucleotide or amino acid residue, the numbering system of Figures 1 and 2 have been maintained.
- 20 The single nucleotide polymorphisms of the present invention which occur within the protein coding sequence may contribute to the phenotype of an organism by affecting protein structure or function. The effect may be neutral, beneficial or detrimental, depending upon the circumstances. Whatever the effect, the identification of such polymorphisms enables for the first time determination of susceptibility to disease, and  
25 new methods of treatment. The single nucleotide polymorphisms of the invention which occur in the non-coding 5' or 3' untranslated regions, may not affect protein sequence, but may exert phenotypic effects by influencing replication, transcription and/or translation. A polymorphism may affect more than one phenotypic trait or may be related to a specific phenotype. In the present invention, polymorphisms in the  
30 corneodesmosin gene are likely to affect the phenotype of an individual with respect to

corneodesmosin-mediated disease, such as inflammatory disease, in particular psoriasis.

5 - The single nucleotide polymorphisms of the corneodesmosin gene, including those of the present invention, are listed in Table 6 where:

- Column 1 designates each single nucleotide polymorphism a reference number.
- Column 2 provides the positional reference of the polymorphism with respect to 10 Figure 1.
- Column 3 indicates position of the SNP with respect to the corneodesmosin coding sequence..
- Column 4 shows the location of the polymorphisms in the gene.
- Column 5 shows the sequence flanking the polymorphism, the polymorphism itself being shown in bold type. For example, the polymorphism at position 6984 is shown as **C/T**, meaning that the variant sequence comprises a T residue, rather 20 than the native C residue.
- Column 6 denotes the standard IUB code.

As discussed above, where a single nucleotide polymorphism of the present invention 25 comprises a nucleotide substitution, the substitution may comprise the replacement of the reference base at a polymorphic site with any other base. Each single nucleotide polymorphism described in Table 6, column 4 represents a preferred embodiment of the invention.

It will be appreciated by those skilled in the art that corneodesmosin gene sequences of the invention may comprise one or more nucleotide substitutions, deletions or insertions in addition to one or more of the single nucleotide polymorphisms of the invention.

5

In a second aspect, fragments of the above polynucleotides are provided, which comprise one or more nucleotide substitutions, insertions or deletions at one or more of the above mentioned positions of Figure 1. Preferably, a fragment may comprise, or even consist of, the polynucleotide sequence of Table 6, column 4. The novelty of 10 a fragment according to the present embodiment may be easily ascertained by comparing the nucleotide sequence of a fragment with sequences catalogued in databases such as GenBank, or by using computer programs such as DNASIS (Hitachi Engineering, Inc.) or Word Search or FASTA of the Genetic Computer Group (Madison, WI).

15

Preferably, the fragments do not encode a full length protein, as is generally the case with the aforementioned polynucleotides, but otherwise satisfy the requirements of the first aspect. Preferred fragments may be 10 to 150 nucleotides in length. More preferably, the fragments are between 5 to 10, 5 to 20, 10 to 20, 20 to 50, or 50 to 100 20 nucleotides in length. For example, the fragments may be 5, 8, 10, 12, 15, 18, 20, 22, 25, 28, 30, or 35 nucleotides in length. The fragments may be useful in a variety of diagnostic, prognostic or therapeutic methods, or may be useful as research tools for example in drug screening.

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In a third aspect of the invention, there is provided non-coding, complementary sequences which hybridise to the corneodesmosin gene sequence. Such "anti-sense" sequences are useful as probes or primers for detecting an allele of a polymorphism of the invention, or in the regulation of the corneodesmosin gene. They may also be used as agents for use in the identification and/or treatment of individuals having or being 30 susceptible to corneodesmosin mediated disease.

The anti-sense sequences of the invention include those which hybridise to an allele of a polymorphism of the invention, and also those which hybridise a region flanking the polymorphic site to enable amplification of an allele of one or more polymorphisms.

5 These sequences may be useful as probes or primers. To be useful as a probe, the anti-sense sequence should bind preferentially one allele of one or more polymorphisms of the present invention and will, preferably, comprise the exact complement of one allele of one or more polymorphisms of the invention. Thus, for example, where the variant comprises a "G" residue at position 7068 of Figure 1, it is preferred that the anti-sense sequence will comprise a "C" residue. Such anti-sense sequences which are capable of specific hybridisation to detect a single base mis-match may be designed according to methods known in the art and described in Maniatis *et al.*, *Molecular Cloning: A Laboratory Manual* 2<sup>nd</sup> Edition (1989), Cold Spring Harbor, NY and Berger *et al.*, *Methods in Enzymology* 152: Guide to Molecular Cloning Techniques (1987) Academic Press Inc. San Diego, CA; Gibbs *et al.*, *Nuc Acids Res.*, 17: 2437 (1989); Kwok *et al.*, *Nuc Acids Res* 18: 999; and Miyada *et al.*, *Methods Enzymol.* 154: 94 (1987). Variation in the sequence of these anti-sense sequence is acceptable for the purposes of the present invention, provided that the ability of the anti-sense sequence to distinguish between alleles of a polymorphism is not compromised. Similarly, variation in the sequence of a primer sequence is acceptable, provided its ability to mediate amplification of a polymorphic site is not compromised. Preferably, a primer sequence will hybridise to the corneodesmosin gene under stringent conditions which are defined below.

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25 In relation to the present invention, "stringent conditions" refers to the washing conditions used in a hybridisation protocol. In general, the washing conditions should be a combination of temperature and salt concentration so that the denaturation temperature is approximately 5 to 20°C below the calculated T<sub>m</sub> of the nucleic acid under study. The T<sub>m</sub> of a nucleic acid probe of 20 bases or less is calculated under

standard conditions (1M NaCl) as [4°C x (G+C) + 2°C x (A+T)], according to Wallace rules for short oligonucleotides. For longer DNA fragments, the nearest neighbour method, which combines solid thermodynamics and experimental data may be used, according to the principles set out in Breslauer *et al.*, *PNAS* 83: 3746-3750 (1986).

5 The optimum salt and temperature conditions for hybridisation may be readily determined in preliminary experiments in which DNA samples immobilised on filters are hybridised to the probe of interest and then washed under conditions of different stringencies. While the conditions for PCR may differ from the standard conditions, the T<sub>m</sub> may be used as a guide for the expected relative stability of the primers. For  
10 short primers of approximately 14 nucleotides, low annealing temperatures of around 44°C to 50°C are used. The temperature may be higher depending upon the base composition of the primer sequence used.

15 The anti-sense polynucleotides of this embodiment may be the full length of the corneodesmosin gene of figure 1, or more preferably may be 5 to 200 nucleotides in length. Preferred polynucleotides are 5 to 10, 10 to 20, 20 to 50, 50 to 100 or 100 to 200 nucleotides in length. Primers, in particular, are typically 10 to 15 nucleotides long, and may occasionally be 16 to 25.

20 In a preferred embodiment, the polynucleotides of the aforementioned aspects of the invention may be in the form of a vector, to enable the *in vitro* or *in vivo* expression of the polynucleotide sequence. The polynucleotides may be operably linked to one or more regulatory elements including a promoter; regions upstream or downstream of a promoter such as enhancers which regulate the activity of the promoter; an origin of replication; appropriate restriction sites to enable cloning of inserts adjacent to the polynucleotide sequence; markers, for example antibiotic resistance genes; ribosome binding sites; RNA splice sites and transcription termination regions; polymerisation sites; or any other element which may facilitate the cloning and/or expression of the polynucleotide sequence. Where two or more polynucleotides of the invention are

introduced into the same vector, each may be controlled by its own regulatory sequences, or all sequences may be controlled by the same regulatory sequences. In the same manner, each sequence may comprise a 3' polyadenylation site. The vectors may be introduced into microbial, yeast or animal DNA, either chromosomal or 5 mitochondrial, or may exist independently as plasmids. Examples of suitable vectors will be known to persons skilled in the art and include pBluescript II, LambdaZap, and pCMV-Script (Stratagene Cloning Systems, La Jolla (USA))

Appropriate regulatory elements, in particular, promoters will usually depend upon the 10 host cell into which the expression vector is to be inserted. Where microbial host cells are used, promoters such as the lactose promoter system, tryptophan (*Trp*) promoter system,  $\beta$ -lactamase promoter system or phage lambda promoter system are suitable. Where yeast cells are used, preferred promoters include alcohol dehydrogenase I or glycolytic promoters. In mammalian host cells, preferred promoters are those derived 15 from immunoglobulin genes, SV40, Adenovirus, Bovine Papilloma virus etc. Suitable promoters for use in various host cells would be readily apparent to a person skilled in the art (See, for example, Current Protocols in Molecular Biology Edited by Ausubel *et al*, published by Wiley).

20 In a fourth aspect of the present invention there is provided a protein or protein fragment comprising an amino acid substitution, deletion or insertion at one or more of positions 18, 130 or 180 of the amino acid sequence of Figure 2. Preferably, the protein or protein fragment is encoded by a polynucleotide according to the first aspect of the invention, and comprises a nucleotide insertion, deletion or substitution at one 25 or more of positions 7164, 10082, 10161, 10162 and 10363 of Figure 1. The corneodesmosin protein or protein fragments of the invention may comprise one or more polymorphisms in addition to one or more of the above-mentioned polymorphisms of Figure 2.

The amino acid sequence exactly as shown in Figure 2 may be referred to as the reference sequence, and is not part of the invention. The amino acid sequence of Figure 2 having an amino acid substitution, deletion or insertion at one or more of the positions indicated above may be referred to as a variant of Figure 2. The reference 5 amino acid at one or more of the above polymorphic sites may be replaced by any other amino acid residue to produce a variant sequence. Amino acid sequences of Figure 2 having one or more of the polymorphisms disclosed in Table 4 are each preferred embodiments of the invention.

- 10 Protein fragments may be functional or non-functional and may be useful in drug screening or gene therapy. Functional fragments may be defined as those which have characteristics of the corneodesmosin protein. The fragments may be at least 10, preferably at least 15, 20, 25 30, 35, 40 or 50 amino acids in length.
- 15 In a fifth aspect of the present invention, there are provided antibodies which react with an antigen of a protein or protein fragment of the fourth aspect. Antibodies can be made by the procedure set forth by standard procedures (*Harlow and Lane, "Antibodies; A Laboratory Manual"* Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1998). Briefly, purified antigen can be injected into an animal in an amount and in intervals sufficient to elicit an immune response. Antibodies can either be purified directly, or spleen cells can be obtained from the animal. The cells are then fused with an immortal cell line and screened for antibody secretion. The 20 antibodies can be used to screen DNA clone libraries for cells secreting the antigen. Those positive clones can then be sequenced as described in, for example, *Kelly et al., Bio/Technology* 10:163-167 (1992) and *Bebbington et al., Bio/Technology* 10:169-175 (1992). Preferably, the antigen being detected and/or used to generate a particular antibody will include proteins or protein fragments according to the fourth aspect.
- 25 In a sixth aspect of the present invention, there is provided host cell comprising a polynucleotide according to any of the aforementioned aspects, for expression of the 30

polynucleotide. The host cell may comprise an expression vector, or naked DNA encoding said polynucleotides. A wide variety of suitable host cells are available, both eukaryotic and prokaryotic. Examples include bacteria such as *E. coli*, yeast, filamentous fungi, insect cells, mammalian cells, preferably immortalised, such as 5 mouse, CHO, HeLa, myeloma or Jurkat cell lines, human and monkey cell lines and derivatives thereof. Such host cells are useful in drug screening systems to identify agents for use in diagnosis or treatment of individuals having, or being susceptible to corneodesmosin mediated disease.

10 The method by which said polynucleotides are introduced into a host cell will usually depend upon the nature of both the vector/DNA and the target cell, and will include those known to a person skilled in the art. Suitable known methods include fusion, conjugation, transfection, transduction, electroporation or injection, as described in Sambrook *et al.*

15 In an seventh aspect of the present invention, there is provided a transgenic non-human animal comprising a polynucleotide according to an aforementioned aspect of the invention. Preferably, the transgenic, non-human animal comprises a polynucleotide according to the first or second aspects. The transgenic animal may be either homozygous or heterozygous for the variant sequence. The animal, and cells derived therefrom, are useful for screening biologically active agents that may modulate corneodesmosin function. Such screening methods are of particular use for determining the specificity and action of potential therapies for corneodesmosin mediated disease, such as psoriasis. The animals are useful as a model to investigate 20 the role of corneodesmosin in normal skin function. Transgenic non-human animals are also useful for the analysis of the single nucleotide polymorphisms and their phenotypic effect.

25 Expression of a polynucleotide of the invention in a transgenic non-human animal is usually achieved by operably linking the polynucleotide to a promoter and/or enhancer

sequence, preferably to produce a vector of the fourth aspect, and introducing this into an embryonic stem cell of a host animal by microinjection techniques (Hogan *et al.*, A Laboratory Manual, Cold Spring harbour and Capecchi *Science* (1989) 244: 1288-1292). Preferably, the construct to be introduced into the animal additionally 5 comprises a) a first homology region with substantial identity to a first corneodesmosin gene sequence; and b) a second homology with substantial identity to a second corneodesmosin gene sequence. The first and second homology regions are of sufficient length for homologous recombination to occur with an endogenous corneodesmosin gene. Those embryonic stem cells comprising the desired 10 polynucleotide sequence may be selected, usually by monitoring expression of a marker gene, and used to generate a non-human transgenic animal. Preferred host animals include mice and other rodents. Further development of such an embryonic stem cell may produce a transgenic animal having cells that are descendant from the embryonic stem cell and thus carry the variant sequence in their genome. Such 15 animals can then be selected and bred to produce animals having the variant sequence in all somatic and germ cells. Such mice can then be bred to homozygosity.

In a preferred embodiment, the transgenic non-human animal may comprise an anti-sense nucleic acid sequence of the third aspect. The expression of an anti-sense 20 sequence in a transgenic non-human animal may be useful in determining the effects of such sequences in treating corneodesmosin-mediated disease, or in neutralising deleterious effects of variant corneodesmosin genes in an animal. Preferably, the host animal will be one which suffers from corneodesmosin mediated disease. The disease 25 may be naturally occurring or artificially introduced.

In some preferred embodiments, for example where the mediated disease has been artificially induced, the transgenic non-human animal will be modulated to no longer expresses the endogenous corneodesmosin gene. Such animals may be referred to as "knock out". In some cases, it may be appropriate to modulate the expression of the 30 endogenous corneodesmosin gene, or express the polynucleotides of the present

invention, in specific tissues. This approach removes viability problems if the expression of a gene is abolished or induced in all tissues. Preferably, the specific tissue would be skin. Where the heterologous gene is human, the animal may be useful in identifying agents which inhibit expression or activity of the variant 5 corneodesmosin sequences of the invention, either *in vivo* or *in vitro*.

In an eighth aspect of the present invention there is provided a method of screening for agents for use in the prognosis, diagnosis or treatment of individuals having, or being susceptible to, corneodesmosin-mediated disease, said method comprising contacting a 10 putative agent with a polynucleotide or protein according to an aforementioned aspect of the present invention, and monitoring the reaction there between. Preferably, the method further comprises contacting a putative agent with a reference polynucleotide or protein of Figure 1 or 2 respectively, and comparing the reaction between (i) the agent and the polynucleotide or protein encoding the reference allele; and (ii) the agent 15 and polynucleotide or protein of the invention. Potential agents are those which react differently with a variant of the invention and a reference allele. It is envisaged that the present method may be carried out by contacting a putative agent with a host cell or transgenic non-human animal comprising a polynucleotide or protein according to the invention. Putative agents will include those known to persons skilled in the art, 20 and include chemical or biological compounds, such as anti-sense polynucleotide sequences, complementary to the coding sequences of the first aspect, or polyclonal or monoclonal antibodies which bind to a product such as a protein or protein fragment of the second aspect. The agents identified in the present method may be useful in determining susceptibility to corneodesmosin-mediated disease, or in the diagnosis, 25 prognosis or treatment of said disease.

In a ninth aspect of the present invention, there is provided a method of diagnosing, or determining susceptibility of a subject to corneodesmosin-mediated disease, said method comprising determining which allele of one or more of the polymorphisms of 30 the invention is present in a subject. The above method may be used in diagnosing or

- determining susceptibility of a subject to any disease in which corneodesmosin is implicated in the pathology, in particular inflammatory disease, such as psoriasis. The method of the ninth aspect may also be used to identify the presence of a combination of single nucleotide polymorphisms in a subject which define a haplotype linked to 5 corneodesmosin mediated disease. The haplotype may be any particular combination of the above single nucleotide polymorphisms, optionally including known polymorphisms. Preferred haplotypes are those shown in Table 10a, the most preferred haplotype being B of Table 10a.
- 10 Any method, including those known to persons skilled in the art, may be used to determine which allele of one or more polymorphisms of the invention is present. Preferably, the method comprises first removing a sample from a subject. More preferably, the method comprises isolating from a sample a polynucleotide or protein to determine therein which allele of one or more polymorphisms of the invention is 15 present.
- Any biological sample comprising cells containing nucleic acid or protein is suitable for this purpose. Examples of suitable samples include whole blood, semen, saliva, tears, buccal, skin or hair. For analysis of cDNA, mRNA or protein, the sample must 20 come from a tissue in which the corneodesmosin gene is expressed, and thus it is preferable to use skin samples.
- In a preferred embodiment, the method for diagnosing, or determining susceptibility of 25 a subject to a corneodesmosin-mediated disease, comprises determining which allele of one or more polymorphisms of the invention is present, in a polynucleotide. Any method for determining alleles in a polynucleotide may be used, including those known to persons skilled in the art. Preferably, the method may comprise the use of anti-sense polynucleotides, as defined above. Such polynucleotides may include sequences which are able to distinguish between alleles of one or more polymorphisms 30 of the invention, by preferential binding, and sequences which hybridise under

stringent conditions to a region either side of a polymorphism of the invention to enable amplification of one or more of the polymorphisms.

Methods of this embodiment include those known to persons skilled in the art, for example direct probing, allele specific hybridisation, PCR methodology including Allele Specific Amplification (ASA), and RFLP.

Determination of an allele of a polymorphism using direct probing involves the use of anti-sense sequences of the third aspect of the invention. These may be prepared synthetically or by nick translation. The anti-sense probes may be suitably labelled using, for example, a radiolabel, enzyme label, fluoro-label, biotin-avidin label for subsequent visualization in, for example, a southern blot procedure. A labelled probe may be reacted with a sample DNA or RNA, and the areas of the DNA or RNA which carry complimentary sequences will hybridise to the probe, and become labelled themselves. The labelled areas may then be visualized, for example by autoradiography.

Allele specific amplification (ASA) discriminates between alleles of a polymorphism on the basis of primers which carry 3' nucleotides specific for a particular polymorphism. Typically, first and second forward primers are provided, wherein the first forward primer hybridises to one allele of a polymorphism of the invention, and the second forward primer comprises a mis-match at the polymorphic site, thus preventing hybridisation. These primers are used in combination with a backward primer, which hybridises to a distal site to enable amplification of the region between a forward primer and the backward primer. As the first forward primer will only bind to a polymorphic site with which it exhibits perfect complementarity, amplification of the region between the forward and backward primers will indicate the presence of a particular allele. The second forward primer having a mis-match at the polymorphic site will not hybridise to the particular allele of a polymorphism, and the absence of a amplification product when this primer is used indicates the absence of the

polymorphism. Preferably, the forward primer will be an anti-sense sequence according to the third aspect of the invention. Preferably, the first forward primer will comprise the complement of a single nucleotide polymorphism of the invention at the 3' most position. The backward primer may hybridise to any suitable portion of the 5 corneodesmosin gene to enable amplification of the intervening region. (see, for example, WO93/22456)

Thus, in a preferred embodiment there is provided a method for diagnosing or determining susceptibility of a subject to corneodesmosin-mediated disease, said 10 method comprising removing a sample from a subject and isolating the nucleic acid therefrom; contacting the sample with either a forward primer which preferentially hybridises to one allele of one or more polymorphisms of the present invention or a forward primer which comprises a mis-match at the polymorphic site and does not hybridise thereto, and a backward primer which hybridises to a distal site; subjecting 15 the nucleic acid sample to amplification; and monitoring for presence of an amplification product which is indicative of the presence of a particular allele of one or more of the polymorphisms of the invention. Preferably, a first reaction is performed using one of the forward primers, and a control reaction is then performed using the other forward primer. It is envisaged that a number alleles of the single 20 nucleotide polymorphisms of the invention may be detected in a single reaction by using multiple primer pairs. Amplification products may then be distinguished by size, using techniques known in the art such as gel electrophoresis, or southern blotting. This method allows the unambiguous identification of individuals homozygous for either allele as well as heterozygous individuals.

25

"RFLP" refers to restriction fragment length polymorphism and is defined as a method of discriminating between two alleles based upon differences in sequence which result in the presence or absence of a restriction enzyme recognition site. In a preferred embodiment of the present aspect there is provided a method for diagnosing or 30 determining susceptibility to corneodesmosin-mediated disease, said method

comprising removing a nucleic acid sample from a subject, and contacting with one or more appropriate restriction enzymes. The size of fragments produced is indicative of which allele of one or more single nucleotide polymorphism according to the invention is present. An allele of a polymorphism of the invention may naturally 5 produce a restriction enzymic site, thus allowing for determination of its presence by analysis of the restriction fragments produced. In some cases, however, an allele of a polymorphism does not create a restriction enzyme site, and one must be artificially introduced. This may be done by using a suitable mis-match primer, according to methods known in the art.

10

The appropriate restriction enzyme, will, of course, be dependent upon the polymorphism and restriction site, and will include those known to persons skilled in the art. Preferred restriction enzymes are listed in Table 3 (ii), column 11, with the expected fragments sizes in columns 7, 8 and 9. Analysis of the digested fragments 15 may be performed using any method in the art, for example gel analysis, or southern blots.

Preferably, the method may first comprise the amplification of a region of the corneodesmosin gene containing one or more of the polymorphic sites of the 20 invention, for example, using PCR techniques. The probes of the present invention may be useful for this purpose.

The above described methods may require amplification of the DNA sample from the subject, and this can be done by techniques known in the art, such as PCR (see *PCR Technology: Principles and Applications for DNA Amplification* (ed. H. A. Erlich, 25 Freeman Press, NY 1992; *PCR Protocols: A Guide to methods and Applications* (eds. Innis *et al.*, Academic press, San Diego, CA 1990); Mattila *et al.*, *Nucleic Acids Res.* 19 4967 (1991); Eckert *et al.*, *PCR Methods and Applications* 117 (1991) and US 30 Patent No 4, 683, 202. Other suitable amplification methods include ligase chain reaction (LCR) (Wu *et al.*, *Genomics* 4 560 (1989); Landegran *et al.*, *Science* 241

1077 (1988)), transcription amplification (Kwoh *et al.*, *Proc Natl Acad Sci USA* **86** 1173 (1989)), self sustained sequence replication (Guatelli *et al.*, *Proc Natl Acad Sci USA* **87** 1874 (1990)) and nucleic acid based sequence amplification (NASBA). The latter two methods both involve isothermal reactions based on isothermal transcription  
5 which produce both single stranded RNA and double stranded DNA as the amplification products, in a ratio of 30 or 100 to 1, respectively.

It may often be desirable to identify the presence of multiple single nucleotide polymorphisms in a sample from a subject. This may be the case in the present  
10 invention where the corneodesmosin gene contains 39 polymorphisms, each of which may be indicative of a different phenotype. For this purpose, nucleic acid arrays may be useful, as described in WO95/11995. The array may contain a number of probes, each designed to identify one or more of the above single nucleotide polymorphisms of the corneodesmosin gene, as described in WO95/11995.  
15

In a further preferred embodiment of the ninth aspect, the method may comprise determining which allele of one or more polymorphisms is present in a protein of the invention. Any method for determining the presence of a particular form, or allele, of a protein is present, may be used. One such method involves the use of antibodies in  
20 diagnosing or determining susceptibility to corneodesmosin mediated disease. The method may comprise removing a sample from a subject, contacting the sample with an antibody to an antigen of a protein or protein fragments according to the second aspect of the present invention, and detecting binding of the antibody to the antigen, wherein binding is indicative of the presence of a particular allele or form of the protein and thus risk to corneodesmosin mediated disease. Tissue samples as  
25 described above are suitable for this method.

The detection of binding of the antibody to the antigen in a sample may be assisted by methods known in the art, such as the use of a secondary antibody which binds to the first antibody, or a ligand. Immunoassays including immunofluorescence assays (IFA)  
30

and enzyme linked immunosorbent assays (ELISA) and immunoblotting may be used to detect the presence of the antigen. For example, where ELISA is used, the method may comprise binding the antibody to a substrate, contacting the bound antibody with the sample containing the antigen, contacting the above with a second antibody bound to a detectable moiety (typically an enzyme such as horse radish peroxidase or alkaline phosphatase), contacting the above with a substrate for the enzyme, and finally observing the colour change which is indicative of the presence of the antigen in the sample.

- 10 In a tenth aspect of the invention, there is provided a method of treating a subject who has been diagnosed as having, or being susceptible to, corneodesmosin mediated disease such as psoriasis. The mode of treatment will depend upon the nature of the polymorphism(s) and the phenotypic effect, and preferably comprises negating the effect of the disease causing polymorphism(s). Where a subject has been diagnosed according to the methods of the invention, treatment to negate the effect of the disease causing polymorphism may include any suitable means. A suitable treatment includes the administration of a polynucleotide sequence which hybridises, preferably under stringent conditions (as defined above), to the corneodesmosin gene. Such polynucleotide sequences may include the anti-sense sequences of the third aspect.
- 15 Alternatively, the treatment may comprise a polynucleotide sequence encoding the corneodesmosin gene or a fragment thereof, and having either a reference or variant allele of a polymorphism of the invention. Preferably, the method comprises:
- 20 (i) determining which allele of one or more polymorphisms of the invention are present; and
- 25 (ii) administering a polynucleotide sequence which hybridises under stringent conditions to the corneodesmosin gene; or a polynucleotide sequence encoding the reference sequence of the corneodesmosin gene or a fragment thereof, or a polynucleotide sequence of the first aspect.
- 30 In an alternative embodiment of this aspect, there is provided the use of a

polynucleotide sequence of the tenth aspect in the manufacture of a medicament for use in the diagnosis and treatment of corneodesmosin mediated disease.

This method of diagnosis and treatment may comprise determining and introducing 5 alleles in the form of a polynucleotide or protein. In the above embodiments, the allele of a polymorphism may be determined using any method, as discussed above. The treatment may be introduced in the form of a protein, or polynucleotide. Any suitable means for introduction of a protein may be used. Introduction of a polynucleotide may use gene therapy methods including those known in the art. In 10 general, a polynucleotide encoding the allele will be introduced into the target cells of a subject, usually in the form of a vector and preferably in the form of a pharmaceutically acceptable carrier. Any suitable delivery vehicle may be used, including viral vectors, such as retroviral vector systems which can package a recombinant genome. The retrovirus could then be used to infect and deliver the 15 polynucleotide to the target cells. Other delivery techniques are also widely available, including the use of adenoviral vectors, adeno-associated vectors, lentiviral vectors, pseudotyped retroviral vectors and pox or vaccinia virus vectors. Liposomes may also be used, including commercially available liposome preparations such as Lipofectin ®, Lipofectamine ®, (GIBCO-BRL, Inc. Gaithersburg, MD), Superfect ® (Qiagen Inc, 20 Hilden, Germany) and Transfectam ® (Promega Biotech Inc, Madison WI).

The polynucleotide or vehicle may be administered parenterally (eg, intravenously), transdermally, by intramuscular injection, topically or the like. As corneodesmosin mediated diseases are usually manifested in the skin, topical administration is preferred. The exact amount of polynucleotide or vehicle to be administered will 25 vary from subject to subject and will depend upon age, weight, general condition, and severity or mechanism of the disorder.

In a further aspect, the present invention provides a kit for the detection in a subject of 30 a single nucleotide polymorphism according to the present invention. Preferably, the

kit will contain polynucleotides according to the aforementioned aspects, most preferably the anti-sense sequences of the third aspect for use as probes or primers; antibodies of the fifth aspect; or restriction enzymes for use in detecting the presence of a polynucleotide, protein or protein fragment of the invention. Preferably, the kit 5 will also comprise means for detection of a reaction, such as nucleotide label detection means, labelled secondary antibodies or size detection means. In yet a further preferred embodiment, the polynucleotides, or antibodies may be fixed to a substrate, for example an array, as described in WO95/11995.

10 The preferred embodiments of each aspect apply to the other aspects of the invention, *mutatis mutandis*.

The present invention will now be described by way of a non-limiting example, with reference to the following figures in which:

15 FIGURE 1 shows the nucleotide sequence of the genomic clone of the corneodesmosin gene, of GenBank Accession No. AC006163.

20 FIGURE 2 shows the amino acid sequence of the corneodesmosin protein and coding sequence therefor.

FIGURE 3 shows the exon and intron structure of the corneodesmosin gene.

ExamplesDetermination of Gene Structure

The mRNA sequence of the corneodesmosin gene (GenBank Accession ID NM\_001264) was used to screen the following public DNA databases: (available through the National Centre for Biotechnology Information website - <http://www.ncbi.nlm.nih.gov/>); NR (Non-Redundant DNA), HTGS (High Throughput Genomic Sequence), and GSS (Genome Survey Sequence). The analysis was performed using the BLASTN algorithm (Altschul, *et al.*, (1990) *J. Mol. Biol.* **215**:403-410). Any genomic sequences containing the corneodesmosin gene were identified by their degree of sequence identity. The gene structure was determined by comparison of the mRNA sequence with the genomic clones. The deduced exon-intron organisation of the corneodesmosin gene is presented in Figure 3.

15 Oligonucleotide primer design for corneodesmosin gene sequencing

5 pairs of oligonucleotide primers (S1F/S1R; S2.1F/S2.1R; S2.2F/S2.2R; S2.3F/S2.3R; S2.4F/S2.4R, S2.5F/S2.5R – Table 1) were designed to amplify exons 1 and 2 of the corneodesmosin gene including 350bp 5' untranslated region (UTR) and 909bp 3' UTR sequences. Oligonucleotide primer sequences were derived from human chromosome 6p21 genomic DNA sequence (GenBank Accession AC006163).

Table 1: Oligonucleotide Primer DNA Sequences.

25

Primer ID	Primer Sequence
S 1F	DCTGGGTCCCGTGGCAAGA
S 1R	DGTCCCTCTCCCGGAGTCTC
S 2.1F	DGGTGAGGGAGGAAGCCAAG

S 2.1R	DGAGCTGACGCTTG GCCAC
S 2.2F	DGCCAACCAATGACA ACTCTTACC
S 2.2R	DGCCTCCACAGAGCTGGAC
S 2.3F	DGGCAAATACTCTCCAGCAACC
S 2.3R	DGGCCTCTCCC ATATGGGA
S 2.4F	DCCAAGGAGAGTTACTCGACAG
S 2.4R	DGGCATATTGGGTGGGTTGAC
S 2.5F	DCATCTGGAAACAGTGGCCAC
S 2.5R	DGTCTTCCTCCTCTGTGGGAG

Corneodesmosin gene amplification

Genomic DNA from a panel of 24 unrelated individuals was amplified using primer pairs S1F/S1R; S2.1F/S2.1R; S2.2F/S2.2R; S2.3F/S2.3R; S2.4F/S2.4R, S2.5F/S2.5R. 5 100ng genomic DNA was amplified by PCR in a total reaction volume of 25µl containing 50mM KCl, 20mM Tris.HCl (pH 8.4), 2mM MgCl<sub>2</sub>, 200µM each dATP, dCTP, dGTP, dTTP, 1µM each oligonucleotide primer and 0.5 units AmpliTaq Gold DNA polymerase (Applied Biosystems). Reactions were thermocycled with an initial 10 denaturation step of 95°C/10mins followed by 35 cycles of 94°C/30secs; T<sub>m</sub> annealing/30secs; 72°C/30secs. A final elongation step of 72°C/10mins completed the amplification. Annealing temperatures (T<sub>m</sub>) for specific primer pairs are presented in Table 2.

15 Table 2: Primer Annealing Temperatures and Amplimer Sizes.

Amplimer	Primer Pairs	Fragment size (bp)	Tm (°C)
1	S1F and S1R	495	63
2.1	S2.1F and S2.1R	610	62
2.2	S2.2F and S2.2R	619	62

2.3	S2.3F and S2.3R	621	63
2.4	S2.4F and S2.4R	532	59
2.5	S2.5F and S2.5R	474	61

Heteroduplex analysis using DHPLC:

5 Oligos were designed to amplify products of between 400-800bp in length from the genomic DNA of 12-25 individuals. Denaturing high-performance liquid chromatography (DHPLC) analysis was performed using the WAVE™ DNA fragment analysis system (Transgenomic) (Kuklin, *et al.*, (1997-98) *Genet Test*. 1(3): 201-6.). The temperature required for successful resolution of heteroduplex molecules  
10 within each PCR product was determined empirically by injecting PCR product at a series of increasing mobile phase temperatures and constructing a fragment specific melting curve. A universal gradient for double stranded DNA was used to determine the appropriate acetonitrile concentration for the heteroduplex identification. For mutation detection, 1-2µl aliquots of the PCR reactions from each of the eleven  
15 individuals were injected onto the WAVE™ column. Mutation detection gradients were for four minutes. Results were graphically visualised using the D-7000 HSM software (Transgenomic).

Direct sequencing of PCR products

20 50-100ng of PCR products were sequenced in both orientations using the DYEnamic ET terminator cycle sequencing premix kit from Amersham. Reactions were fractionated on ABI 377 automated sequencers using standard procedures. Chromatographic traces were analysed using the SEQUENCHER programme (Gene Codes, USA), to identify SNP  
25 positions.

Detection of Variant Alleles - Assay design for genotyping

The fragment sequence containing the polymorphism was analysed for the creation or deletion of a naturally occurring restriction enzyme recognition site in response to variation in the nucleotide sequence. If the polymorphism did not result in any changes in restriction enzyme recognition sites then the sequence was interrogated with the Primer Design Mismatch Program™. This is an adaptation of the program described by Davidow LS ((1992) *Comput Appl Biosci* 8:193-194).

10      Detection of Polymorphisms in 24 Population Controls

The application of the approach outlined above resulted in the identification of 39 SNPs. These are described in Table 3, in which:

- Column 1 designates each single nucleotide polymorphism a reference number.
- Column 2 provides the positional reference of the polymorphism with respect to Figure 1, together with details of the polymorphism itself. For example, the reference "C6948T" indicates a substitution of the nucleotide "C" for nucleotide "T" at position 6984 of Figure 1.
- Column 3 of (i) provides the corresponding positional references with respect to the coding sequence of the corneodesmosin gene.
- Column 4 of (i) indicates the region of the gene which the polymorphism occurs.
- Column 5 of (i) shows the sequence flanking the polymorphism, the polymorphism itself being shown in bold type. The single nucleotide polymorphisms are defined using standard IUB code.

- Columns 3 and 4 of (ii) show primer sequences which may be used to amplify a region of the corneodesmosin gene to enable detection of the single nucleotide polymorphism by using restriction enzyme analysis. The amplified product size is shown in column 5 of (ii).

5

- Columns 6 to 9 of (ii) list the restriction enzymes used to digest the amplified product, and the sizes of fragments generated by the reference, variant and heterozygous sequences respectively.

10 RFLP or ASA assays were developed for all of these SNPs and the corresponding primers along with amplification product and digestion fragment sizes are also given in Table 3. Of these 39 SNPs, 9 give rise to amino acid changes. These are shown in Table 4.

15 Additional Corneodesmosin Polymorphisms

20 In a subsequent experiment, DNAs from 96 individuals comprising 24 type IA psoriatics, 24 type IB psoriatics, 24 type II psoriatics and an additional 24 healthy controls, were sequenced as described above using primers designed to cover the remainder of the Corneodesmosin gene (see Table 5a)

The sequencing reactions were carried out with 50-100ng of PCR products sequenced in both orientations using the DYEnamic ET terminator cycle sequencing premix kit from Amersham according to the following protocol:

25

The PCR products were Exo/Sap treated and desalted using p10 columns, prior to setting up the sequencing reactions in a thermowell plate including:

200-400ng PCR Product

30 1µl primer @ 10pmolml<sup>-1</sup>

8 $\mu$ l ET Termination mix

H<sub>2</sub>O to 20 $\mu$ l

The plates were sealed with an MJ Research Microseal film and then vortexed to mix  
5 samples, followed by a spin to ensure reaction is at the bottom of the wells.

PCR was carried out according to the following protocol:

No Predenaturation  
95 °C for 30sec  
10 50 °C for 15 sec  
60 °C for 1 min  
for 40 cycles and then hold at 10 °C until ready to purify.

After removing the plate from the thermocycler, the products were purified by ethanol  
15 precipitation. To each well we added 2 $\mu$ l 7.5M ammonium acetate followed by 80 $\mu$ l  
100% ethanol and incubated at room temperature for 10 minutes before spinning at  
4000rpm for 1 hour at room temperature. The supernatant was discarded and the  
pellet washed with 70% ethanol before centrifugation for a further 30 minutes. The  
supernatant was discarded and remaining ethanol removed gently by pipetting using  
20 p10 tips before allowing the pellets to air dry.

The samples were then resuspended in 10 $\mu$ l MegaBACE Loading Buffer (Molecular  
Dynamics) and transferred to a Robbins plate prior to loading onto the MegaBACE.  
Reactions were fractionated on a Molecular Dynamics MegaBACE capillary  
25 sequencer using standard procedures. Chromatographic traces were analysed using  
the SEQUENCHER programme (Gene Codes, USA), to identify SNP positions.

A total of 28 novel SNPs were identified (additional to those given in the example above). For reference, these are SNPs 6-18 and 53-67 in Table 5b. A combined list of Corneodesmosin SNPs is given in Table 6.

5      Corneodesmosin gene association with psoriasis

A total of 21 SNPs (see Table 7) were genotyped in 147 families identified through a proband with psoriasis (a total of 499 individuals, of whom 233 were affected). The genotyping was carried out using a variety of methods (single base extension using the 10 Snapshot kit from Amersham Pharmacia Biotech, Pyrosequencing (Ahmadian A *et al.*, Anal Biochem 2000 280:103-10), or direct sequencing) as given in Table 7. All these methods used established methodologies that are provided by the equipment manufacturers and/or are well known to those skilled in the art.

15      Linkage Disequilibrium

The extent of linkage disequilibrium (LD) between markers was calculated using genotype data from 199 unrelated, unaffected individuals and is expressed as correlation coefficients in Table 8. This analysis shows that there is extensive linkage 20 disequilibrium between many of the Corneodesmosin polymorphisms.

Single point association

Single point associations between each SNP and psoriasis affected status were 25 calculated using the TRANSMIT program (Clayton D, MRC Biostatistics Unit, Cambridge) - see Table 9. Highly significant associations were observed between SNPs 19, 21, 23, 24, 26, 28, 30, 33, 34, 37, 38 and psoriasis. The single SNP showing the most significant association with psoriasis that has been previously reported is 30 SNP 33 (Tazi Ahnini R *et al.*, Hum. Mol. Genet. 1999: 8 pp1135-40; Allen MH *et al.*, Lancet 1999: 353 pp1589-90).

This study has identified 9 SNPs, (19, 21, 24, 26, 28, 30, 34, 37 and 38) which show global chi-squared values greater than that seen for SNP 33, and are therefore more powerfully predictive of affected status.

5

#### Haplotype analysis

A total of 19 SNPs were used for haplotype analysis (SNPs at positions 29 and 32 were excluded due to low information content). Three common haplotypes were 10 identified (Table 10). Of the three common haplotypes, haplotype B is significantly associated with psoriasis. The alleles are coded alphabetically (table 10b) such that the nucleotide first in the alphabet is given coded as 1, and the other nucleotide is coded as 2. Thus A is always 1, T is always 2, and G or C are coded depending on the other nucleotide. For example, in SNP No. 1, which is a C to T substitution, the 15 presence of the C allele is coded as 1 and the presence of the T allele is coded as 2 (see Table 10b). In Table 10a, this means that haplotypes A and B have C residues, and haplotype C has a T residue at this position. For an A to C substitution, the A allele will be coded as 1, and the C allele as 2. In a C to G substitution, the C allele will be 1 and the G allele 2.

20

#### Construction of Corneodesmosin Gene Targeting Vector

As the genetic data pointed strongly to an involvement of the Corneodesmosin gene in the pathophysiology of psoriasis, we decided to engineer mouse strains in which the 25 mouse orthologue of the corneodesmosin gene is knocked out by homologous recombination using a vector construct designed to remove exon 2 of the Corneodesmosin gene.

Murine Corneodesmosin genomic clones were isolated from a mouse large insert PAC 30 library, using mouse Corneodesmosin cDNA sequence as a probe by standard

techniques. The isolated murine Corneodesmosin genomic clones were then restriction mapped in the region of the Corneodesmosin gene using small oligonucleotide probes and standard techniques. The murine genomic locus was partially sequenced to enable the design of homologous arms to clone into the targeting vector. The murine 5 Corneodesmosin gene is a two-exon gene. A 4 kb 5' homologous arm and a 1 kb 3' homologous arm where amplified by PCR and the fragment cloned into the targeting vector. The position of these arms was chosen to functionally disrupt the Corneodesmosin gene by deleting the majority of the coding sequence. A targeting vector was prepared where the deleted Corneodesmosin sequence was replaced with 10 non-homologous sequences composed of an endogenous gene expression reporter (an in frame fusion with lacZ) upstream of a selection cassette composed of a self promoted neomycin phosphotransferase (neo) gene in the same orientation as the Corneodesmosin gene.

15 Transfection and Analysis of Embryonal Stem Cells

Embryonal stem cells (Evans MJ & Kaufman MH Nature 1981 292:154-6) were cultured on a neomycin resistant embryonal fibroblast feeder layer grown in Dulbecco's Modified Eagles medium supplemented with 20% Fetal Calf Serum, 10% 20 new-born calf serum, 2 mM glutamine, non-essential amino acids, 100 $\mu$ M 2-mercaptoethanol and 500 u/ml leukemia inhibitory factor. Medium was changed daily and ES cells were subcultured every three days. 5.times.10.sup.6 ES cells were transfected with 5  $\mu$ g of linearized plasmid by electroporation (25  $\mu$ F capacitance and 400 Volts). 24 hours following electroporation the transfected cells were cultured for 9 25 days in medium containing 200  $\mu$ g/ml neomycin. Clones were picked into 96 well plates, replicated and expanded before being screened by PCR to identify clones in which homologous recombination had occurred between the endogenous Corneodesmosin gene and the targeting construct. From 96 picked clones 45 targets were identified. These clones where expanded to allow replicas to be frozen and

sufficient high quality DNA to be prepared for Southern blot confirmation of the targeting event using external 5' and 3' probes, all using standard procedures (Russ *et al.* Nature 404:95-99).

5      Generation of Corneodesmosin Deficient Mice

C57BL/6 female and male mice were mated and blastocysts were isolated at 3.5 days of gestation. 10-12 cells from Clone 7 (described in Example 2) were injected per blastocyst and 7-8 blastocysts were implanted in the uterus of a pseudopregnant F1 10 female. Five chimeric pups were born of which one male was 100% agouti (indicating cells descended from the targeted clone). This male chimera was mated with female and MF1 and 129 mice, and germline transmission was determined by the agouti coat color and by PCR genotyping respectively.

15      Corneodesmosin knock-out mouse as a model of corneodesmosin-mediated disease

Mice heterozygous for the Corneodesmosin knockout are superficially normal. Staining for expression of the lacZ reporter gene fused to the Corneodesmosin promoter in the knockout construct shows clear expression in desquamating skin. We 20 then genotyped surviving offspring from intercrosses of heterozygous knockout mice on an outbred genetic background in an attempt to isolate mice homozygous for the knockout.

From 44 surviving progeny we identified:

- 25      17 wild type  
          27 heterozygotes  
          0 homozygous mutant.

Statistical analysis of these data indicate that the ratio of wild type:heterozygous animals conforms to a 1:2 ratio consistent with a homozygous lethal phenotype (Chi square = 0.557).

- 5 In keeping with this analysis, two pups found dead 24-48 hours after birth were homozygous mutant. Together these data indicate the Corneodesmosin deficiency in mice is lethal with pups dying soon after birth, most likely through dehydration as a result of failure to establish a permeability barrier in the skin.
- 10 We conclude from this that altering the activity of Corneodesmosin (e.g. by modulating expression or altering its proteolytic processing) will be useful in developing models of disease in which epithelial integrity is increased (e.g. psoriasis) or decreased (e.g. dermatitis), and for testing novel agents for the alleviation of Corneodesmosin mediated disease.

Table-3(j) S Gene SNPs with location and assay details

SNP	SNP	Corneodesmosin		Location	Flanking Sequence
		nt position	nt position		
1	C6984T	-115	5' UTR	CTCCCCGCCA CACCAACTTC CCCYGGCA CCCACCCCT CCACCTCTCC	AATGTCCAGOTCTGGCATAA AGGACCGGG TGTCCTCGAG CTGCCATCAG
2	A7068G	-31	5' UTR	TCTGGCATAA AGGACCGGG TGTCCTYGA GCTGDCATCAG TCAAGGAGGCC	
3	C7077T	-22	5' UTR	CTGCCCAGTCAG TCAAGGAGGCC GCTCGGGCA	
4	C7107T	9	5' UTR	GGCGTGTGGGTGGGCACATGWTGGCACT GCTGCTGCCTGCTCTC	
5	A7164T	66	Coding Sequence	CTAAAGAGCAT TGGCACCTC TCAGACCCYT GTAAGGACCCCACGGGTATC	ACCTCCCTAACGACCCCTGCGYTACTGGGAAGGGTG
6	C1039T	137	Coding Sequence	CAGTAGCTAC AGTGGCTCCA GCAYTCTGG CAGCTCATTTCAGTGCCA	
7	C1082T	180	Coding Sequence	GAGGAGCAGC TGTCACTCGG GAARCA GGGCTCTCACTCG GGAAAGCAGCA	
8	C10134T	206	Coding Sequence	GAAGCAGGGGCTCTCACTCG GGAA(G)AGCA GCTCTCAATTGAGCAGC	
9	G10344A	442	Coding Sequence		
10	10353(AAG)ins	481	Coding Sequence		
11	A10516G	614	Coding Sequence	CTGGACAAAGGCTTCCCTCTCCCTCTCCCTCTCCCTCTCCCTCTCCCTCTCC	CTGGACAAAGGCTTCCCTCTCCCTCTCCCTCTCCCTCTCCCTCTCCCTCTCC
12	C10521T	619	Coding Sequence	GGAGGGCCCCA TCGTCTCGCA CTCYGGCCCC TAGATCCCCA GCTCCCACTC	GGAGGGCCCCA TCGTCTCGCA CTCYGGCCCC TAGATCCCCA GCTCCCACTC
13	T10624C	722	Coding Sequence	GCTCCCACTCCTGTCAGGG GGTCAAGAGRC CTGTTGGTGT GTGGGTGAGAC	GCTCCCACTCCTGTCAGGG GGTCAAGAGRC CTGTTGGTGT GTGGGTGAGAC
14	G10669A	767	Coding Sequence	CCTACAGTA GGTTAAATC TAYCCTGTTGG OCTACTTCAC CAAGAGAAC	CCTACAGTA GGTTAAATC TAYCCTGTTGG OCTACTTCAC CAAGAGAAC
15	T10873C	971	Coding Sequence	AGCCAGTGGCAGGTTCCCTC GGCCATTGCR TTCAAGCCAG TGGGGACTGG	AGCCAGTGGCAGGTTCCCTC GGCCATTGCR TTCAAGCCAG TGGGGACTGG
16	G11020A	1118	Coding Sequence	CTCCCTCCAGTTCTCGAGTC OCCAGAORGTT CTAGCAATTTCAGCA GTC	CTCCCTCCAGTTCTCGAGTC OCCAGAORGTT CTAGCAATTTCAGCA GTC
17	A11117G	1215	Coding Sequence	CCCAGCAGTTCTAGCAATTTCAGCA GGGTCAACCCTAACGATCCCTG	CCCAGCAGTTCTAGCAATTTCAGCA GGGTCAACCCTAACGATCCCTG
18	T11138G	1236	Coding Sequence	CTAGCAATTTCAGCA GGGTCAACCCTAACGATCCCTGCGGCAGTGTCT	CTAGCAATTTCAGCA GGGTCAACCCTAACGATCCCTGCGGCAGTGTCT
19	G11142T	1240	Coding Sequence	CTAGCAATTTCAGCA GGGTCAACCCTAACGATCCCTGCGGCAGTGTCT	CTAGCAATTTCAGCA GGGTCAACCCTAACGATCCCTGCGGCAGTGTCT
20	C11145T	1243	Coding Sequence	GGCAAGAGCTC CAGTTGCCAA TCSAGTGGCA AAATCATCCTTGT	GGCAAGAGCTC CAGTTGCCAA TCSAGTGGCA AAATCATCCTTGT
21	G11233C	1331	Coding Sequence	TCGAGTGGCA AAATCATCCTTGT	TCGAGTGGCA AAATCATCCTTGT
22	T11260C	1368	Coding Sequence	TTCCTACCCC AAGGAGAGTT ACTCRACAG CCATAAGTCA ACTGTTGTGT	TTCCTACCCC AAGGAGAGTT ACTCRACAG CCATAAGTCA ACTGTTGTGT
23	G11495A	1593	Coding Sequence	GAGAGTTACTCTGAGCAGTTCAGTTCAGTTCAGTTCAGTTCAGTTCAGTTC	GAGAGTTACTCTGAGCAGTTCAGTTCAGTTCAGTTCAGTTCAGTTCAGTTC
24	11536(AAG)ins	1603	3' UTR		
25	G11576T	1674	3' UTR	TACACTATATCCCATATGGGAGAAGKCCAGTGCCTCAAGGCATAGGGTTAGC	CCAAAAGAAGTGTGTTCTGCTTCTCYACTACCCCTAAGGTTGCAAGACTCTC
26	T11641C	1739	3' UTR	AGTGGGTTCTGCTTCTACTAACCCYAAAGGTTGCAAGACTCTCTCTTATCA	CCCTTACAATTCCCTCTACTGTGKAATTTGTCATTGAGTAACACCC
27	T11649C	1747	3' UTR	GGTCAATTGAGTAACACCCATCASC TTCTAACCTGGAAACCCCTGAA	GTAAATGCTCTCAGAGCACCTCTGAYGCCTGAAGAAGTTATCCTTCCCTC
28	T11808G	1806	3' UTR	AAAAAGTGGCAGCCAACTTCAGTTCAGTTCAGTTCAGTTCAGTTCAGTTC	AAACAGTGGCAGCCAACTTCAGTTCAGTTCAGTTCAGTTCAGTTCAGTTC
29	C11839G	1937	3' UTR	TTATOTOAAGCCCCCTTCTGTCAGTGCCTGCCCAGATGATTCCTG	CAACCCAACTTCCCTCTGTCAGTGCCTGCCCAGATGATTCCTG
30	C11885T	1983	3' UTR	TICCTGGCCATTTCAGTGCCTGTCAGTGCCTGTCAGTGCCTGTCAGTGCCTG	TTTATGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG
31	C11977T	2075	3' UTR	GACACTGGGGACCCCTAGGGTGRAGCAGGCTCCCTGCTGCTGCTGCTGCTG	GGTCAAGAC TTTTGCCTTCTGTCAGTGCCTGCTGCTGCTGCTGCTGCTG
32	T12018G	2116	3' UTR	CTCTCTGTCAGTGCCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG	TCCTCTGAGAGCTGGGGTGTGGTGTGGTGTGGTGTGGTGTGGTGTGGTGT
33	T12136C	2234	3' UTR	CCTCTGTCAGTGCCTGTCAGTGCCTGTCAGTGCCTGTCAGTGCCTGTCAG	CCTCTGTCAGTGCCTGTCAGTGCCTGTCAGTGCCTGTCAGTGCCTGTCAG
34	C12149T	2247	3' UTR	TCATTTCTCTTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCT	TCATTTCT
35	G12198A	2295	3' UTR		
36	G12283A	2381	3' UTR		
37	T12318C	2416	3' UTR		
38	C12345T	2443	3' UTR		
39	C12373A	2474	3' UTR		

Table 3(ii)

**Table 4- Amino Acid Polymorphisms**

SNP	POSITION	LOCATION	VARIANT 1	VARIANT 2	Effect on amino acid side chain
5	A7164T	EXON 1	MET	LEU	Conservative
6	C10039T	EXON 2	PRO	PRO	Neutral
7	C10082T	EXON 2	LEU	SER	Hydrophobic - Hydrophilic
8	C10108T	EXON 2	GLY	GLY	Neutral
9	G10344A	EXON 2	SER	ASN	Conservative
10	10363 (AAG)ins	EXON 2	SER insertion	SER deletion	SER insertion/deletion
11	A10516G	EXON 2	GLN	GLN	Neutral
12	C10521T	EXON 2	SER	PHE	Hydrophilic - Hydrophobic
13	T10624C	EXON 2	SER	SER	Neutral
14	G10669A	EXON 2	ARG	ARG	Neutral
15	T10873C	EXON 2	TYR	TYR	Neutral
16	G11020A	EXON 2	ALA	ALA	Neutral
17	A11117G	EXON 2	SER	GLY	Hydrophilic - Hydrophobic
18	T11138G	EXON 2	SER	ALA	Hydrophilic - Hydrophobic
19	G11142T	EXON 2	GLY	VAL	Conservative
20	C11145T	EXON 2	SER	LEU	Hydrophilic - Hydrophobic
21	G11233C	EXON 2	SER	SER	Neutral
22	T11260C	EXON 2	CYS	CYS	Neutral
23	G11495A	EXON 2	ASP	ASN	Hydrophilic charged - Hydrophilic neutral

Table 5

Primer Name	Primer Sequence Forward	Primer Sequence Reverse
SEEKINI_8	CAGTGAGCTGAGACCCGTG	CTGGTACCACTGTTGTCAG
SEEKINI_8	CAGTGAGCTGAGACCCGTG	CTGGTACCACTGTTGTCAG
SEEKINI_8	CAGTGAGCTGAGACCCGTG	CTGGTACCACTGTTGTCAG
SEEKINI_6	GACTCCTCAGAGCCTCAG	GTAGCTACTGAAGCCGCTG
SEEKPROM3	CCTAGATCAAGAGCCCCAG	ACAGCAGGAGACTCGAGG
SEEKPROM2	CCTCAGATGCTTCATGAATGG	GTGAAGTCAGCCGAATAGC

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Table 6 cont'd

Table 7

SNP	AC006163	nt position*	SNP	chemistry	Frequency	
					allele 1	allele 2
1	6,984bp	44884 CDSN6984		PSQ	69.6	30.4
2	7,058bp	44968 CDSN7068		PSQ	60.6	39.2
19	10,059bp	47939 PS SEEKIN1_6 C565T	Sequenced		55	45
21	10,108bp	48008 CDSN C10098T	Sequenced	not available	not available	
22	10,344bp	48244 CDSN G10343A	Sequenced	not available	not available	
23	10,363bp (ins)	48262 CDSN 10363 AA G ins	Sequenced	not available	not available	
24	10,516bp	48416 CDSN x2 2A10516G	PSQ	47.8	52.2	
25	10,521bp	48421 CDSN x2 2C10521T	PSQ	20.5	79.5	
26	10,624bp	48524 CDSN x2 T10614C	SNaPshot	48.9	51.1	
27	10,669bp	48569 CDSN x2 2G10669A	SNaPshot	-	85.7	14.3
28	10,873bp	48773 CDSN T10873C	SNaPshot	32.3	67.7	
29	11,020bp	48920 SEEKIN1_3 G27A	PSQ	43.8	56.2	
30	11,117bp	49017 SEEKIN1_3 A124G	PSQ	38.8	1.2	
31	11,135bp	49038 SEEKIN1_3 T145G	PSQ	82.8	17.2	
32	11,142bp	49042 SEEKIN1_3 G149T	PSQ	100	0	
33	11,145bp	49045 SEEKIN1_3 C152T	PSQ	64.3	35.7	
34	11,233bp	49133 SEEKIN1_3 G241C	PSQ	47.8	52.2	
35	11,280bp	49160 SEEKIN1_3 T268C	PSQ	78.9	21.1	
36	11,495bp	49395 SEEK1in3 G503A	SNaPshot	68.7	31.3	
37	11,505bp (ins)	49404-49407 SEEK1in3_511INS	SNaPshot	43.4	56.6	
38	11,575bp	49479 CDSN G11576T	SNaPshot	32.5	67.5	

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Table 8

Table 9

SNP Number	Position	SNP Type	Frequency (allele1)	Transmissions	Chi-squared (bootstrap)	P value	Number of Transmissions		
							Allele 1 observed	Allele 1 expected	Allele 2 observed
1	44884	Promotor	0.79	117	3.92	0.043	235	226	53
2	44968	Promotor	0.69	98	1.14	0.213	157	152	67
19	47939	Silent	0.79	133	11.43	0.002	260	244	50
21	48008	leu>ser	0.74	125	10.7	0	238	221	60
22	48244	ser>asn	0.2	132	3.44	0.061	55	64	259
23	48262	ins/del (ser)	0.82	112	9.28	0	231	219	33
24	48416	Silent	0.59	126	16.03	0	203	180	99
25	48421	ser>phe	0.18	120	1.43	0.18	44	49	232
26	48524	silent	0.43	140	22.93	0	113	143	229
27	48569	silent	0.13	139	5.97	0.025	35	45	305
28	48773	silent	0.58	142	36.51	0	152	188	194
29	48920	silent	0.47	26	0.88	0.283	23	26	33
30	49017	ser>gly	0.96	131	11.16	0	291	299	21
31	49038	ser>ala	0.13	135	4.55	0.051	34	42	290
32	49042	gly>val	1						282
33	49045	ser>leu	0.59	132	9.69	0	211	193	111
34	49133	ser>leu	0.43	133	11	0.002	115	135	203
35	49160	silent	0.33	102	0.74	0.381	77	81	168
36	49395	silent	0.22	140	6.47	0.02	61	74	281
37	49404	ins/del	0.58	139	18.32	0	223	197	113
38	49479	3'UTR	0.44	144	34.99	0	194	158	192

Table 10a

SNP Number	Haplotype		
	A	B	C
1	1	1	2
2	1	1	2
19	1	1	2
21	1	1	2
22	2	2	1
23	1	1	1
24	1	1	2
25	1	2	2
26	2	2	1
27	2	2	2
28	2	2	1
30	1	1	1
31	2	2	2
33	1	1	2
34	2	2	1
35	2	2	1
36	2	2	1
37	1	1	2
38	1	1	2

Table 10b

Key	Code	
	1	2
A/T	A	T
A/G	A	G
A/C	A	C
C/G	G	C
G/T	G	T
C/T	C	T

Claims

1. A recombinant or isolated polynucleotide comprising a nucleic acid sequence encoding the corneodesmosin gene of Figure 1, wherein said corneodesmosin gene comprises a nucleotide substitution, deletion or insertion at one or more of positions 5 6984, 7068, 7077, 7107, 7164, 8884, 8906, 8931, 9538, 9607, 9608, 9647, 9667, 9745, 9761, 9926, 9952, 9968, 10082, 10161, 10162, 10363, 11567, 11641, 11649, 11808, 11839, 11885, 11977, 12018, 12136, 12149, 12198, 12283, 12318, 12345, 12373, 10 12901, 13001, 13020, 13108, 13117, 13178, 13224, 13316, 13365, 13562, 13605, 13670, 13859, 13889 and 13914 of Figure 1.
2. A recombinant or isolated polynucleotide comprising a nucleic acid sequence encoding a fragment of the corneodesmosin gene of Figure 1, wherein said fragment comprises a nucleotide substitution, deletion or insertion according to claim 1. 15
3. A recombinant or isolated polynucleotide comprising a nucleic acid sequence which hybridises under stringent conditions to the corneodesmosin gene.
- 20 4. A recombinant or isolated polynucleotide according to claim 3, wherein said nucleic acid preferentially hybridises to one allele of one or more of the polymorphisms of claim 1.
- 25 5. A recombinant or isolated polynucleotide according to claim 3 wherein the nucleic acid sequence hybridises under stringent conditions to a region of the corneodesmosin gene flanking one or more of the polymorphisms of claim 1.
6. A vector comprising a polynucleotide according to any one of claims 1 to 5.
- 30 7. A host cell comprising a polynucleotide or vector according to any one of

claims 1 to 6.

8. A protein comprising the amino acid sequence of Figure 2 and having an amino acid substitution, deletion or insertion at one or more of positions 18, 130 or 5 180 of Figure 2, or a fragment thereof.

9. An antibody or antibody fragment which binds to a protein or protein fragment according to claim 8.

10 10. A transgenic non-human animal comprising a polynucleotide sequence according to any one of claims 1 to 6.

11. Use of a transgenic non-human animal according to claim 10 in screening for agents for use in diagnosis or treatment of individuals having, or being susceptible to, 15 corneodesmosin mediated disease.

12. A method of screening for an agent for use in the prognosis, diagnosis or treatment of individuals having or being susceptible to corneodesmosin mediated disease, said method comprising contacting a putative agent and with a polynucleotide or protein according to claims 1 to 5, or monitoring the reaction there between. 20

13. A method of screening for an agent according to claim 13, further comprising contacting a putative agent with a polynucleotide or protein of figures 1 or 2 respectively; and comprising the reaction between  
25 (i) the agent and polynucleotide or protein of claim 1 to 5, or 8; and  
(ii) the agent and a polynucleotide or protein of Figures 1 or 2.

14. A method of diagnosing for, or determining susceptibility to, corneodesmosin mediated disease, comprising determining which allele of one or more polymorphisms 30 of claim 1 or 7 are present in a subject.

15. A method according to claim 14, said method comprising determining in a protein or protein fragment which allele of one or more polymorphisms of claim 8 are present in a subject.

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16. A method for diagnosing and treating corneodesmosin mediated disease in a subject, comprising

- (i) determining which allele of one or more polymorphisms of the invention are present; and
- 10 (ii) administering a polynucleotide sequence which hybridises under stringent conditions to the corneodesmosin gene; or a polynucleotide sequence encoding the reference sequence of the corneodesmosin gene or a fragment thereof, or a polynucleotide sequence of the first aspect.

15 17. Use of a polynucleotide sequence according to claim 16 in the manufacture of a medicament for use in the diagnosis and treatment of corneodesmosin mediated disease.

18. A kit for use in diagnosis of an individual having, or being susceptible to, 20 corneodesmosin mediated disease, said kit comprising an agent for detection of a polynucleotide, protein or protein fragment according to any one of claims 1 to 5 or 7 or 8, together with a key correlating the alleles of one or more polymorphisms with presence of, or susceptibility to, corneodesmosin mediated disease.

25 19. A kit according to claim 18 wherein the agent comprises a polynucleotide according to claim 3, an antibody according to claim 9 or restriction enzymes for digestion of a polynucleotide according to claim 1 or 2.

**FIG. 1**Genomic Sequence of Corneodesmosin Gene

6701 TGGAGGGCAG ATGGAGAGAC AGGCCAAGCC ACGGTAGGCA GGAGAGTTAA  
6751 GGAGCCAGGC AGCTGGTCC CGTGGCAAGA GTGGCCGCC CAGAGTGGT  
6801 GGCGGTGGG CAGAGCGCCT GGTTCCGGGT TAGGCAATGA GGAGCCGGG  
6851 CCAGGCCTGT CAGGTGGCAG GATCGTTAGA GCCCCGTGGC CATGGTACC  
6901 CCACACTGCA GCCACTGCTG CTGCTGAGTA GGCAGATGCA CCGGGCTGAT  
6951 TACGACGGTC CCTCCGGGCA CAUCACTTC CCCCGGGCA CCCACCCG  
7001 GGACCTCTTC TCCTCTGCC ACASTGACTC TGCCCAGGG AAATGTCAGG  
7051 TCTGGATAAA AGGACCCCAGG TCTCTCCAG TGCCCATCAG TCAGGAGGCC  
7101 GTGCAGCCCG AGATGGGCTC GTCTCGGGCA CCCTGGATCG GGCGTGTGGG  
7151 TGGGCACGGG ATGATGGCAC TGCTGCTGGC TGGTCTCCTC CTGCCAGGTA  
7201 GGAGGCCTGGG GCCCCTGGGA ACAGGAGGGGA GGCGGGAGGG AGACTCCGGG  
7251 AGAGGACCCA GCGAAGGGGA CGGGCAGGGG CTCTGGAATC TGCCTTTGA  
7301 GTCTGGGGT TGCTCCTCAC TGTATGGTCG CCTCAGGTAA GTTTCTAAA  
7351 CTPCCTGAGC CCCAGTTCT GAAATTCTGA AGTGGGGTTA ATGACACCTA  
7401 CCTCTAGTCT GTGTGTCTCA AATTAAATAA TGTATGTGAT ATGTACTTTG  
7451 GAAATTCTAG AGGTTATAT AAATGGTGGT GGTGATTTT ATTATGGGAG  
7501 CACTACAAGA TAATGATTGG ACATTTAATA GTAATAATAT CATTGGAGA  
7551 GCCTTTTAT ATGCTAGACT CTGTTTAAG CACATTTGGA TTATATATTA  
7601 GAACTTTAT TTTTATTTT TTTGTGAGAT GGAGTCCCAC TCTGTCTCCA  
7651 AGGCTGGAGT GCAGTGGCGT AATCTCGGCT CACTGCAACT TCCACCTCTC  
7701 AGGTTCAAGC GACTCTCATG CCTCAGCCTC TAGAGTAGCT GGGACAACAG  
7751 GTGCCCATCA CCACACCTGG CTAATTCT TTTTTTGTA TTTTTAGTAG  
7801 AACACAGGGTT TTACCAATTG GGTCAAGCTG GTCTGAACT CCTGACTCAA  
7851 GTGATCCGCT CGCCTCGGCC TCCCAAGGTG CTGGGATTAC AGGCATGAGC  
7901 CACCACACCC GGCCTATATT AGCACTTTG ATCATFACAA GAACGGTATG

7951 AAAAGAGATT TGCTATTCC ACTCTACAGA TGAGGACACT GAGGCTCGGA  
 8001 GAGGTAGGA AACTAGCTCA AAATCATGCA TTAGAAGGCA GCAAAGCCAA  
 8051 GATTCAACC CCAGGCCAGG CAACCCCTGG ACCTGTGTTG TTGACCACCG  
 8101 GGTACTTATA GCCCTTGAGG AATTCTGCG ACCTTCCCCT GGTCTAGTGG  
 8151 GTGGTTGGTG TCTGAGGGAA TAGCGAAAGA GAGAGGCAAT GCATGGTGGA  
 8201 TTCGTGCAGA GGACTGAAGG GAATTGGCAC AGCTGGGTT CGGCCTGGAG  
 8251 GTGCATGCAG AGAATTCTT TCTGAGGAGA GAACAGGGAC ATCACAGAGG  
 8301 ATGGCAGTCT GGTGTTGGT GGAGGGATCA GGATGAGTGG CAGTAATAAT  
 8351 TCATAATATA TAATGCTTTA CACTTTCTAA AACATCTGGC CGCACATGAT  
 8401 AGCTTGTGCC TGTAATCCA ACACCTCAGG AGGCCAAGGC AGGTGAATCG  
 8451 CCTGAGGTCA GGAGTTCAAG ACCAGCCTGG CCAAGATGGT GAAACCCCT  
 8501 CTCTACTAAA AATACAAAAAA ATTAGCTGGG TGTGGTGGCG GGCACCTGTG  
 8551 GTCCCAGCTA CTTGGGAGGC TGAGGCAGGA GAATCGCTTG CACCAAGGAG  
 8601 GCAGAGGTTA CAGTGAGCTG AGACCGTGTT ATTGCACTTT AGCCTGGCA  
 8651 ACAAGAAACT CCATCTCACA AAAAAAAAAA AAAAAAAA AAAGAAGAAA  
 8701 AAACCTCCAG GTGGATGATC TCATTTAGTT TTCTTCATAG TAATGCTGTG  
 8751 GGAAGGCAGG GAAAATTGG CCCCTCTGAA TGTATAAACT AAAGCTCAGA  
 8801 GAGGTTCACT AACTTGCTAG TATGTGGCTC TGTTGTAAAC ACGTGGGACC  
 8851 TGGAGGGGCT AGGGAAGGCA GAAGGAACGC AGGTGAAAGA GTCATGGAGG  
 8901 AACCATGGGG TAAGTTGGC CTGGGGTTT GAGCAAAGGA AAGGAAAGAT  
 8951 AAGGAAAGAT GTGGCTCCAC ATCCCTGAGG GAAGTCAAGG CAGCAGAACT  
 9001 CAGATGAGGG GCTGGACAGA GGCAAGGTGTG CTCAGAGAGG GAAGCTGATT  
 9051 GTGGCCAGGA GCCTCGGAGG TTCGTGGGT TTCTGCCTGG TTCCCTGGC  
 9101 TGGGCCAGCG AGAGCAGGGC TGGCTCAGGG TGCGGTGTCC TGACACACTG  
 9151 GTACCAGCAG GTTCTGAAGC AACAGGTAGT GACCCACAT CCTGGCCCCC  
 9201 ACCCAGCTTT ACTGGCATGG CCAGTGC~~T~~GA GATAGGAAAT AGGGTTCCA

9251 TTCCCTGACCC CAGCCTGGGC TCTCACGAAG AAGCTGGTGA CCAAATCTTA  
 9301 GTCCTCGAGT GCCCTTTCCT TTATTTCA~~G~~C CCCCTGCCC CCAGCTTG~~T~~  
 9351 CT~~T~~TTTCCAG TGTCTCCTTC TATATGTGTC TCCACTTCTC AGCCCTCCAT  
 9401 TGTTTG~~C~~CCT TTTGTCTTCT TCCCTG~~G~~GT CCCACTGTCT GGCCCAGGAT  
 9451 TTTTCCC~~T~~A AGAATTTACG CCTGGACTCC TCAGAGCCTC AGTTTCCCCA  
 9501 ATTCTCTGTC TCTTCAGGGT CCTTTCTTT AGACCTATT~~T~~ GTTCCTGCC~~C~~  
 9551 CTTCTCCATT CCCTCTTCTT TTTAAAAAAA ATTTAATT~~A~~ AAAAACAAAA  
 9601 TACAGATGGG GTCTATGTTG CCCAGGCTGG TCTTGAAC~~T~~C TGGGGCC~~C~~AT  
 9651 GCAATCCTCC CACCTCAGCC TCCC~~A~~AGTG CTGGGATTAC CGGC~~G~~TGAGC  
 9701 CACTGTGCC~~C~~ AGCCCCCTCT TATATTCAAT CTATT~~C~~OTT~~T~~ GAGGTCACTC  
 9751 ACTTTGGCAC GTATTTCT ATTTTCTGG TTGGTGT~~T~~TG CCCACC~~T~~TC  
 9801 CCAAACAAAG AAATGCCTT ATT~~C~~GGCAC C~~T~~CAATATCC TTAGAGACA  
 9851 ATAGCCAGTT CTTCC~~T~~CC~~T~~ TCTCC~~A~~CCCC TAAACTCTCC CTGC~~G~~CTCTG  
 9901 CTTGGGAGAA ACCCGAGAGG CCGATTACTG AGATAAGGCA GAAAGGTGAG  
 9951 GGAGGAAGCC AAGCCTCTT GGCC~~C~~TACT AACCA~~T~~GCT TT~~C~~CTCCACA  
 10001 GGGACCTTGG CTAAGAGCAT TGGCAC~~CT~~TC TCAGACCC~~T~~ GTAAGGACCC  
 10051 CACCGTATC ACCTCCCC~~T~~A ACGACCC~~T~~G CCTCA~~T~~GGG AAGGGTGACT  
 10101 CCAGCGGCTT CAGTAGCTAC AGTGGCT~~C~~CA GCAGTTCTGG CAGCTCCATT  
 10151 TCCAGTG~~C~~CA GAAGCTCTGG TGGTGGCTCC AGTGGTAGCT CCAGCGGATC  
 10201 CAGCATTGCC CAGGGTGGTT CTGCAGGATC TTTAAGGCC GGAACGGGGT  
 10251 ATTCCCAGGT CAGCTACTCC TCCGGATCTG GCTCTAGTCT ACAAGGTGCA  
 10301 TCCGGTTCC~~T~~ CCCAGCTGGG GAGCAGCAGC TCTCA~~T~~CGG GAAGCAGCGG  
 10351 CTCTCACTCG GGAAGCAGCA GCTCTCATTC GAGCAGCAGC AGCAGCTT~~C~~  
 10401 AGTTCAGCAG CAGCAGCTTC CAAGTAGGGA ATGGCTCTGC TCTGCCAAC~~C~~  
 10451 AATGACA~~A~~CT CTTACCGCGG AATACTAAAC CCTTCCCAGC CTGGACAAAG  
 10501 CTCTTCC~~T~~CT TCCC~~A~~ACCT CTGGGGTATC CAGCAGTGGC CAAAGCGTCA

10551 GCTCCAACCA GCGTCCCTGT AGTTGGACA TCCCCGACTC TCCCTGCAGT  
 10601 CGAGGGCCCCA TCGTCTCGCA CTCTGGCCCC TACATCCCCA GCTCCCACTC  
 10651 TGTGTCAGGG GGTCAAGAGGC CTGTGGTGGT GGTGGTGGAC CAGCACGGTT  
 10701 CTGGTGCCCCC TGGAGTGGTT CAAGGTCCCC CCTGTAGCAA TGGTGGCCTT  
 10751 CCAGGCAAGC CCTGTCCCCC AATCACCTCT GTAGACAAAT CCTATGGTGG  
 10801 CTACGAGGTG GTGGGTGGCT CCTCTGACAG TTATCTGGTT CCAGGCATGA  
 10851 CCTACAGTAA GGGTAAAATC TATCCTGTGG GCTACTTCAC CAAAGAGAAC  
 10901 CCTGTGAAAG GCTCTCCAGG GGTCCCTTCC TTTGCAGCTG GGCCCCCAT  
 10951 CTCTGAGGGC AAATACTTCT CCAGCAACCC CATCATCCCC AGCCAGTCGG  
 11001 CAGCTTCCTC GGCCATTGCG TTCCAGCCAG TGGGGACTGG TGGGCTCCAG  
 11051 CTCTGTGGAG GCGGCTCCAC GGGCTCCAAG GGACCCCTGCT CTCCCTCCAG  
 11101 TTCTCGAGTC CCCAGCAGTT CTAGCATTTC CAGCAGCTCC GGTCACCCCT  
 11151 ACCATCCCTG CGGCAGTGCT TCCCAGAGCC CCTGCTCCCC ACCAGGCACC  
 11201 GGCTCCTTCA GCAGCAGCTC CAGTTCCAA TCGAGTGGCA AAATCATCCT  
 11251 TCAGCCTTGT GGCAGCAAGT CCAGCTCTTC TGGTCACCCCT TGCATGTCTG  
 11301 TCTCCTCCTT GACACTGACT GGGGGCCCCG ATGGCTCTCC CCATCCTGAT  
 11351 CCCTCCGCTG GTGCCAAGCC CTGTGGCTCC AGCAGTGCTG GAAAGATCCC  
 11401 CTGCCGCTCC ATCCGGGATA TCCTAGCCCA AGTGAAGCCT CTGGGGCCCC  
 11451 AGCTAGCTGA CCCTGAAGTT TTCCTACCCCA AAGGAGAGTT ACTCGACAGT  
 11501 CCATAAGTCA ACTGTTGTGT GTGTGGCATGC CTTGGCCACA AACAAAGCACA  
 11551 TACACTATAT CCCATATGGG AGAAGGCCAG TGCCCCAGCCA TAGGGTTAGG  
 11601 TCACTTTCCC TCCCTYCCCAA AAGAGTGGTT CTGCTTTCTC TACTACCCCA  
 11651 AGGTTGCAAGA CTCCTCTGTCA TCACCCCTTC CTCCTTCCTC TTCTCAAAAT  
 11701 GCTAGATTGA AAGCTCTCTC CTTGATTCTC TCCTACTGTT TAAATTECCCA  
 11751 TTCCACCGACA GTGCCCTCA GCCAGATCAC CACCCCTTAC AATTCCCTCT  
 11801 ACTGTGTGA AATGGTCCAA TGAGTAACAC CCCCATCACC TTCTCAACTG

FIG. 1 CONT'D

11851 GCAAACCCCT GAAATGCTCT CAGASCAGCT CTGACGCCCTG AAGAACGTAT  
 11901 ACCTTGCCTCT TCGGCTTTCAG CAATAAAGG AAAGTCAAAC CATCACTGG  
 11951 AACACTGGC CACTTTTCAC TGACCTCTCT TCGACATCTA GTGAAACCCAC  
 12001 GAAATATGCC ACTGGGCTTT CGCTCCCAAT TCCACCCCCAC CCTCCATTAC  
 12051 AGACCTCACCCACGGGCTCT AGATCACCGT CCCCAACACA CCCATTGCCCT  
 12101 CTCAAGGCCC TTAATCTCAGG CCCATTCTCTG GGGCAATTCC CTCAGTGCCC  
 12151 AGATCATTCG CTGGGTGAGG GAGACACTGG GGACACCTICA GAGGTTGGAG  
 12201 CAGGTCCTCTGCTGTCCTG CTCCTGGAG AGATGGCTCA GTAAACHTG  
 12251 GGGACTTACGT GCAGACCTTT TGCCCTCTTG GAGTCCTGGG TCTCTCTCTGA  
 12301 CACTCTGGGT GCTGCTCTTC CTACGCCCTCT AGAGGTCTCT GTGTCCTCTCA  
 12351 TTTCCTCTICA AAAGGGGGCT GTGTTCTCTG TCTACCTCTC ACCTCCCTCC  
 12401 ACAGAGGAGG AAGACAATAA ATAAATTGTTG AACATGAAACG AGAGATPGCG  
 12451 TGCCCTCCCA GATGCTTCCG CCATTCCCT CCTCTCTCAT TGCTCCAGGA  
 12501 AATCCATTCI CTCCTCCATTC CTGATTCACCG GTGGGGTCCC CCTTCGGCTT  
 12551 ATTTAGGGGG CTCAGTGTCTT TCTCTCCGTC CCCTTECCCTC CCCCTCCCCAC  
 12601 CCAAACCTCC TTTGTTCCAG CATTAGCATT CCTCACCTTC TAGATGCCAT  
 12651 CGCTGTCGGG AGTCAATGAGE CTCGATTTCC TGGGTTCTG GGACACCTGG  
 12701 AAGCTTGGGA AGGCTGGGAC ACAACAACTC CAACCAAGTT CCTGTCAGCT  
 12751 GAGTAGGAGG CCAAGTTCGGC GTTGTTCCTG GAGCTGGGGG TGGAGAGAGT  
 12801 AAAGGACTGA GAGGATGGGA CGGGGGAGG GAGTGCAGCG AAGCAGGGTG  
 12851 ACTCACTGGC CTAGATCAAG AGGCCAGCC TGTGGCAGAA CAGAGCTGCC  
 12901 ACTGGTCCTCT CCATCTTCAC ACTCCCTGCT CTGCTGGGT CCACACTGAG  
 12951 AGTGTGAGCA ACATGGCTCT CAGGTGAGGG CTGAGAAGGC AGAGTCCCCG  
 13001 AGTGGAAAG AGGAGTCGCT TCCACTGGAG AAGAGAGAGA AAGTGGAGTG  
 13051 TGTGGTGGGG TCCATGCGAC TTAACTCCTG AGACAGGGAG GGAGAGGCTG  
 13101 AGCCGGAGGA AGTCCCCGCA TCCCAACGGAG GGCAGAGTGG ATTGTGCTTG

13151 TCCCCTGTAGG AGGCCCAACCC CCCACCCCCAG CCCACGTCTC AGAGGCTCTG  
 13201 CTTGGGTCCA AAGGAATTCA CCCCTAGTGT AGGACTTAAC CCATTCGCTC  
 13251 CTATCAGGGT GGTGGTGTGTTGGTCCCTGAAT TTAGAAGTGTGTTGAAATCGAA  
 13301 AGTGTGGAAT CACCAAAAAT GTATTACATT GACCAAGAAAG GGATTCGATC  
 13351 ACCCTTGCTC CAGCACTCG CCTCTGATCT GCAGCCAAATG GCAGGAATCG  
 13401 ACCTCCCTCAC ATCTCTCATC AATGGAAATT GCAGGGAGAG AACGGCTCTC  
 13451 CAIAGTGCTT TCCCTGGACT CTCTTGCTGT CCTCCAAATTAATAGCTTGT  
 13501 CTAAACGCA TGCAGTCAT CGAAAAAGGC CTGTGGGCTC CATCCACTGG  
 13551 CAGTTCTGGA CAGGAGCTCT TCACCTCTCC AGTGGTTAACG CCACOACGGG  
 13601 CAGGTGGGCA CGACACAGCA CTAGAATCAC CGTAGAGCTC ATGTTAGAC  
 13651 CTTGGGCAGC CAGGGAAACC TACTCCCTGG GCCTCCCGGA AGCCATGGAG  
 13701 AGAACAAAGG CATTCCATTT TTATAATAAA AATGGAAAC AAATTTAAAA  
 13751 GCCAACAAAC TGTAAATGAA TCTCTACATT CTCACTGGCC AGCTTCAACA  
 13801 AGGATCCCAG CTTCAACAAG GATCAAGGCC TGGCCATTTC ACAGGAGCAT  
 13851 TTAAACGCTC TCTCTACTG TTACTGAA ATAGGACTTT TCTCCGAAGG  
 13901 TTCTTAAATAC TCTGTGGCAC ATCTGACAC CAGTAGGAGG CAGAATGATG  
 13951 TCTTCAACCC CAACACCATC AAAGATGTC ACATGCTAA CGGTGGAACG  
 14001 TAGGAATTAG GTACATGGC AAAGGAAAT TAAGGTCCA GATGGGATTA  
 14051 AGGTTGGTAT TCGGCTGACT TCACAGAGAT TATCATGGAT TATTCAGGTG  
 14101 GGTCCAGTGT AGTCACCGAGG TCCCTTAATG TGGACATGGG AGGCAGAAGA  
 14151 GGAAGTCTGA GIGATACAGT GTAAGAAATG GCTGATTTG GCTTTGGAGA  
 14201 TGGAGGAAGG GGACCATGAG CCAAGAACAGGGATGCCG CTAGAAGGTG  
 14251 AAAAACGAGG GAAGGGATT TTCCTCTGAG CCCCCCCAGAA ACAATCAGAG  
 14301 CCCTGGTCACT ACCTTTATTTC TAATCCACTG AGACCTGTTT TAGACTTCTG  
 14351 ATCTCGAAAA CTGTAAAGTA ATAATCCAT GTTGTGTAA GCGATTCGGT

UTR Sequence

Exon Sequence

SNP

## FIG. 2

### CORNEODESMOSIN (af030130.em\_hum1)

ATGGGCTCGTCTCGGGCACCCCTGGATGGGGCGTGTGGTGGGCACGGGATGATGGCACTG  
 15 TACCCGAGCAGAGCCCCTGGGACCTACCCCGCACACCCACCCGTGCCCTACTACCGTGAC  
 M G S S R A P W M G R V G G H G M M A L -  
 L

CTGCTGGCTGGTCTCCTCCTGCCAGGGACCTTGGCTAAGAGCATTGGCACCTTCTCAGAC  
 75 GACGACCGACCAGAGGAGGACGGTCCCTGGAACCGATTCTCGTAACCGTGGAAAGAGTCTG  
 L L A G L L P G T L A K S I G T F S D -  
 T

CCCTGTAAGGACCCCACCGGTATCACCTCCCCFAACGACCCCTGCCCTCACTGGGAAGGGT  
 135 GGGACATTCCTGGGGTGCATAGTGGAGGGATTGCTGGGGACGGAGTGACCCCTCCCA  
 P C K D P T R I T S P N D P C L T G K G -  
 S

GACTCCAGCGGCTTCAGTAGCTACAGTGGCTCCAGCAGTTCTGGCAGCTCCATTCCAGT  
 195 CTGAGGTCGCCGAAGTCATCGATGTCACCGAGGTCGTCAAGACCGTCGAGGTAAAGGTCA  
 D S S G F S S Y S G S S S S G S S I S S -  
 T

GCCAGAACGCTCTGGTGGCTCCAGTGGTAGCTCCAGCGGATCCAGCATTGCCAGGGT  
 255 CGGTCTCGAGACCACCACCGAGGTACCATCGAGGTCGCCCTAGGTCGTAAACGGGTCCCA  
 A R S S G G G S S G S S S S G S S I A Q G -  
 T

GGTTCTGCAGGATCTTTAAGCCAGGAACGGGTATTCCCAGGTCAAGCTACTCCTCCGGA  
 314 CCAAGACGTCCTAGAAAATTGGTCCTGCCCTAAGGGTCCAGTCATGAGGAGGCCT  
 G S A G S F K P G T G Y S Q V S Y S S G -  
 T

TCTGGCTCTAGTCTACAAGGTGCATCCGGTTCCCTCCAGCTGGGGAGCAGCAGCTCTCAC  
 375 AGACCGAGATCAGATGTTCCACGTAGGCCAAGGAGGGTCGACCCCTCGTCGAGAGTG  
 S G S S L Q G A S G S S Q L G S S S S H -

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FIG. 2 CONT'D

AAGCCCTGTCCCCAATCACCTCTGTAGACAAATCCTATGGTGGCTACGAGGTGGTGGGT  
 855 -----+-----+-----+-----+-----+-----+-----+-----+----- 914  
 TTCGGGACAGGGGTTAGTGGAGACATCTGTTAGGATACCACCGATGCTCCACCACCA  
 K P C P P I T S V D K S Y G G Y E V V G -  
  
 C  
 GGCTCCTCTGACAGTTATCTGGTCCAGGCATGACCTACAGTAAGGGTAAAATCTATCCT  
 915 -----+-----+-----+-----+-----+-----+-----+-----+----- 974  
 CCGAGGAGACTGTCAATAGACCAAGGTCCGTACTGGATGTCATTCCCATTAGATAGGA  
 G S S D S Y L V P G M T Y S K G K I Y P -  
  
 GTGGGCTACTTCACCAAAGAGAACCCGTGAAAGGCTCTCCAGGGTCCCTTCCTTGCA  
 975 -----+-----+-----+-----+-----+-----+-----+-----+----- 1034  
 CACCCGATGAAGTGGTTCTCTGGGACACTTCCGAGAGGTCCCCAGGAAAGGAAACGT  
 V G Y F T K E N P V K G S P G V P S F A -  
  
 GCTGGGCCCTCATCTCTGAGGGAAATACTTCTCCAGCAACCCATCATCCCCAGCCAG  
 1035 -----+-----+-----+-----+-----+-----+-----+-----+----- 1094  
 CGACCCGGGGTAGAGACTCCCGTTATGAAGAGGTCTGGGTAGTAGGGTCGGTC  
 A G P P I S E G K Y F S S N P I I P S Q -  
  
 A  
 TCGGCAGCTCCTCGGCCATTCGCGTTCAGCCAGTGGGACTGGTGGGTCCAGCTCTGT  
 1095 -----+-----+-----+-----+-----+-----+-----+-----+----- 1154  
 AGCCGTCAAAGGAGCCGGTAACGCAAGGTCTGACCCCTGACCACCCAGGTCGAGACA  
 S A A S S A I A F Q P V G T G G V Q L C -  
  
 GGAGGCCTCCACGGCTCCAAGGGACCCGTGCTCTCCCTCCAGTTCTCGAGTCCCCAGC  
 1155 -----+-----+-----+-----+-----+-----+-----+-----+----- 1214  
 CCTCCGCCGAGGTGCCGGAGGTTCCCTGGGACGAGAGGGAGGTCAAGAGCTCAGGGTCC  
 G G G S T G S K G P C S P S S S R V P S -  
  
 G T T T  
 AGTTCTAGCATTCCAGCAGCGCCGGTTACCCCTACCATCCCTGCGGCAGTGCTTCCCAG  
 1215 -----+-----+-----+-----+-----+-----+-----+-----+----- 1274  
 TCAAGATCGTAAAGGTCTCGCGGCCAAGTGGGATGGTAGGGACGCCGTACGAAGGGTC  
 S S S I S S S A G S P Y H P C G S A S Q -  
 S V L

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1275 C  
 AGCCCCTGCTCCCCACCAGGCACCGGCTCCTCAGCAGCAGCTCCAGTCCAAATCGAGT  
 1334 TCGGGGACGAGGGGTGGTCCGTGGCCGAGGAAGTCGTCGAGGTCAAGGGTAGCTCA  
 S P C S P P G T G S F S S S S S S Q S S -  
  
 1335 C  
 GGCAAAATCATCCTTCAGCCTTGTCAGCAAGTCCAGCTCTCTGGTCACCCCTGCATG  
 1394 CCGTTTTAGTAGGAAGTCGAAACACCGTCGTTCAGGTCGAGAAGACCAGTGGAACGTAC  
 G K I I L Q P C G S K S S S S G H P C M -  
  
 1395 TCTGTCTCCTCCCTGACACTGACTGGGGGCCCGATGGCTCTCCCCATCCGTGATCCCTCC  
 1454 AGACAGAGGAGGAAGTGTGACTGACCCCCGGGGTACCGAGAGGGTAGGACTAGGGAGG  
 S V S S L T L T G G P D G S P H P D P S -  
  
 1455 GCTGGTGCCAAGCCCTGTGGCTCCAGCAGTGCTGGAAAGATCCCTGCCGTCCATCCGG  
 1514 CGACCACGGTTGGGACACCGAGGTGTCACGACCTTCTAGGGGACGGCGAGGTAGGCC  
 A G A K P C G S S S A G K I P C R S I R -  
  
 1515 GATATCCTAGCCCAAGTGAAGCCTCTGGGGCCCAGCTAGCTGACCCCTGAAGTTTCCTA  
 1574 CTATAGGATCGGGTTCACTTCGGAGACCCGGGGTCACTGGACTTCAAAGGAT  
 D I L A Q V K P L G P Q L A D P E V F L -  
  
 1575 A  
 CCCCAAGGAGAGTTACTCGACAGTCCATAA  
 1604 GGGGTTCTCTCAATGAGCTGTCAGGTATT  
 P Q G E L L D S P \* -  
 N

FIG. 2<sub>CONT'D</sub>

